RECYCLE OF NITROXYL-CONTAINING STREAMS AT LOW **TEMPERATURE**

I claim the benefit under Title 35, United States Code, § 120 to U.S. Provisional Application Number 60/222,595, filed August 2, 2000, entitled RECYCLE OF NITROXYL-CONTAINING STREAMS AT LOW TEMPERATURE.

BACKGROUND OF THE INVENTION

1. Field of the Invention

The present invention relates to the inhibition of polymerization and polymer growth of ethylenically unsaturated monomers by means of the addition thereto of at least one stable nitroxide free radical compound.

2. **Description of Related Art**

Many ethylenically unsaturated monomers undesirably polymerize at various stages of their manufacture, processing, handling, storage, and use. Polymerization, such as thermal polymerization, during their purification results in the loss of the monomer, i.e., a lower yield, and an increase in the viscosity of any tars that may be produced. The processing and handling of the higher viscosity tars then requires higher temperature and work (energy cost) to remove residual monomer.

Polymerization can also result in equipment fouling, especially in the case of production of acrylic monomers. Such polymerization causes loss in production efficiency owing to the deposition of polymer in or on the equipment being used. These deposits must be removed from time to time, leading to additional loss in production of the monomer.

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A wide variety of compounds has been proposed and used for inhibiting uncontrolled and undesired polymerization of ethylenically unsaturated monomers.

However, many of these compounds have not been fully satisfactory.

There are several mechanisms by which polymerization inhibitors work. One mode of action for polymerization inhibitors is for the inhibiting species to combine with the propagating polymer chain such that the polymerization of that polymer chain stops, i.e., a termination reaction. If such an inhibitor-terminated polymer chain is capable of participating in a dynamic equilibrium between a dormant species (the inhibitor-terminated chain) and an active polymer chain, it would be considered a "living" or quasiliving polymer. For example, Ivan, Macromol. Symp. 88:201-215 (1994) describes quasiliving polymerization as a polymerization in which "... only a portion of chain ends are active (propagating) and these are in equilibria with inactive (dormant, nonpropagating) chains..." Shigemoto et al., Macromol. Rapid Commun. 17:347-351 (1996) state, "Well-defined polymers can be prepared by controlled / "living" radical polymerization in the presence of relatively stable radicals. These systems employ the principle of dynamic equilibration between dormant species and growing radicals via reversible homolytic cleavage of a covalent bond in dormant species." Further, Greszta et al., Macromolecules 29:7661-7670 (1996) state, "The reversible homolytic cleavage of dormant species can be accomplished by either thermal, photochemical, or catalytic activation. The most successful approaches are as follows: homolytic cleavage of alkoxyamines and dithiocarbamates, use of various organometallic species, and catalyzed atom transfer radical polymerization." Such a

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"living" polymer is capable of increasing in molecular weight (growing) through its reaction with additional monomer units of the same or different types of polymerizable monomers.

The method by which this "living" polymer grows is termed the "living" polymerization mechanism, and is depicted below.

M-Inh ---->
$$M^* + *Inh$$
 (1)

$$M^* + *Inh ----> M-Inh$$
 (2)

$$M^* + M' - M^*$$
 (3)

$$M-M'* + *Inh ----> M-M'-Inh$$
 (4)

Reactions (1) and (2) depict the dynamic equilibrium, with (2) being the termination reaction. Reaction (3) depicts growth of the polymer chain. Reaction (4) depicts retermination of the growing polymer chain with the inhibiting species. The amount of growth over any period of time is dependent on the relative rate at which (2) occurs versus (3), as long as (1) occurs to some extent. The faster (2) is relative to (3), the more time is needed for significant growth of the polymer. Under the conditions in which inhibitors are normally used, the concentration of the inhibiting species should be sufficiently high to cause reaction (2) to be much faster than reaction (3), otherwise it would not be an effective inhibiting system for commercial use. However, we have realized that even at an effective inhibiting amount of the inhibitor, growth can still occur, given sufficient time and temperature.

There are at least two scenarios in which "living" polymer can remain in a monomer purification train for an excessive amount of time.

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First, the use of tar recycle can significantly increase the amount of time that the "living" polymer can remain in the purification train. To recycle unused inhibitor that is left in the purification stream after removal of the monomer, a portion of the residual (or tar) stream is added to a feed stream earlier in the purification train. This residual stream typically contains inhibitor, small amounts of monomer, impurities in the monomer stream that have been concentrated by the purification process, and polymer formed during the production and purification process. Recycling this polymer will allow it time to grow if it is "living" polymer and the conditions of the purification train allow the "living" polymerization mechanism to occur. If this polymer grows via the "living" polymerization mechanism, excessive polymerization would cause loss in product yield, increased waste residues from the process, and potential plugging of equipment due to excessively high molecular weight polymer in the purification stream.

Second, occasionally, conditions in the plant/purification process can result in the formation of polymer within the purification train that is not dissolved by the monomer stream. If this polymer is caught in a dead space, or if it attaches to the metal on the inside of the equipment, it will not be washed out of the system. Thus, the polymer will remain within the system indefinitely (potentially for two or more years). If this polymer grows via the "living" polymerization mechanism, it could coat the inside of the equipment, causing inefficient separation of the monomer stream components and/or insufficient or inefficient heat transfer to enable purification. Such a situation would cause loss in product yield and could potentially cause an unscheduled shut-down of the plant in order to clean out the undissolved polymer in

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the equipment. Such a shut-down results in loss of monomer production and additional expense to clean out and dispose of the undissolved polymer.

It is significant that there has been no indication that previously used inhibitors would lead to the formation of "living" polymer when used as polymerization inhibitors. However, a newly utilized class of inhibitors, the stable nitroxyl radicals, is known to allow this "living" polymerization mechanism to occur. These nitroxyl radicals are highly efficient polymerization inhibitors under normal use, providing better performance than most other inhibitors on the market, but their incapacity to prevent "living" polymerization has hindered their full utilization. Accordingly, there is a need for improved techniques that can be employed in a purification train whereby nitroxyl radicals can be used to full advantage to prevent the polymer growth that occurs via a "living" polymerization mechanism.

Nitroxyl radicals are known to facilitate polymerization via a "living" free radical process to give polymers of narrow polydispersity.

Georges et al., Macromolecules 26(11):2987-2988 (1993) synthesized narrow molecular weight resins by a free-radical polymerization process with polydispersities comparable to those that can be obtained by anionic polymerization processes and below the theoretical limiting polydispersity of 1.5 for a conventional free-radical polymerization process. The process comprised heating a mixture of monomer(s), free-radical initiator, and a stable free radical, e.g., 2,2,6,6-tetramethyl-1-piperidinyloxy (TEMPO).

Hawker *et al.*, *Macromolecules 29(16)*:5245-5254 (1996) prepared and evaluated a variety of initiating systems for the preparation of macromolecules by nitroxide-mediated "living" free radical procedures. The systems were divided into two classes, unimolecular initiators in which alkylated TEMPO derivatives dissociate to provide both the initiating radical and the stable radical, and bimolecular systems in which a traditional free radical initiator, such as BPO or AIBN, is used in conjunction with TEMPO. For the unimolecular initiators it was found that an α -methyl group is essential for "living" character, while a variety of substituents could be placed on the phenyl ring or the β -carbon atom without affecting the efficiency of the unimolecular initiator. It was found that the rate of polymerization is approximately the same for both the unimolecular and corresponding bimolecular systems; however, the unimolecular initiators afforded better control over molecular weight and polydispersity.

Hindered nitroxyl compounds are known to be very active inhibitors of free radical polymerizations of unsaturated monomers such as styrene, acrylic acid, methacrylic acid, and the like. Nitrophenols, nitrosophenols, phenylenediamines (PDA's), hydroxylamines, quinones and hydroquinones are also known to have a similar capacity.

U.S. Patent Number 3,163,677 discloses a process for the preparation of N,N,O-trisubstituted hydroxylamines and N,N-disubstituted nitroxides of the formulae:

$$\begin{bmatrix} R_1 \\ R_2 - C \\ R_3 \end{bmatrix}_2 N - O - C - R_2 \qquad \text{and} \qquad \begin{bmatrix} R_1 \\ R_2 - C \\ R_3 \end{bmatrix}_2 N - O *$$

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wherein R₁, R₂, and R₃ are each an alkyl radical having 1 to 15 carbon atoms. (As used herein, the designation N-O* denotes a stable free radical wherein the asterisk is an unpaired electron.) The N,N,O-trisubstituted hydroxylamines can be used to make the N,N-disubstituted nitroxides, which are stable free radicals and are said to be useful as polymerization inhibitors.

U.S. Patent Number 3,334,103 discloses that nitroxides can be prepared from the corresponding heterocyclic amine wherein the nitrogen atom of the nitroxide group is attached to other than a tertiary carbon of an aliphatic group (i.e., the nitrogen atom forms a part of a heterocyclic nucleus). These nitroxides are said to have useful properties similar to those described for the N,N-disubstituted nitroxides of U.S. Patent Number 3,163,677.

U.S. Patent Number 3,372,182 discloses that a great variety of N,N-disubstituted, stable, free radical nitroxides not otherwise readily available can be prepared by a simple and convenient process that comprises pyrolyzing in an inert reaction medium virtually any hydroxylamine that is susceptible to cleavage of the O-C bond, e.g., tri-t-butylhydroxylamine.

U.S. Patent Number 3,422,144 discloses stable, free radical nitroxides of the formula:

wherein R is selected from the group consisting of tertiary alkyl, aryl, alkaryl, haloaryl, carboxyaryl, alkoxyaryl, alkylthioaryl, pyridyl, and dialkylaminoaryl, and R' is tertiary

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alkyl. These nitroxides are said to be useful as traps for reactive free radicals both in the counting of free radicals and for inhibiting oxidation and free radical polymerization.

U.S. Patent Number 3,494,930 discloses free radicals of the nitroxide type for use as initiators of free radical reactions, collectors of free radicals, polymerization inhibitors or antioxidants. They are constituted by nitrogenous bicyclic compounds in which one of the bridges comprises solely the nitroxide radical group and, in particular, by aza-9-bicyclo (3,3,1) nonanone-3-oxyl-9, and by aza-9-bicyclo (3,3,1) nonane oxyl-9.

U.S. Patent Number 3,873,564 discloses compounds and a method for assaying enzymes by adding to a medium containing an enzyme a stable free radical compound having a stable free radical functionality which, when subjected to an enzyme-catalyzed reaction, changes the environment of the free radical functionality. By following the change in the electron spin resonance spectrum as affected by the change in environment, the type of enzyme and the activity of the enzyme can be determined. The compounds found useful are normally stable nitroxide radicals with an enzyme labile functionality. Other compounds include two cyclic nitroxide containing rings joined by a chain having an enzyme labile functionality.

U.S. Patent Number 3,959,395 discloses a process for recovery and re-use of dinitrophenols employed as inhibitors of polymerization in the distillation and purification of styrene. The styrene still residues commonly referred to as tar or tars are treated with an aqueous hydroxide at a controlled pH, the phases are separated, the aqueous phase is treated with an acid and an organic solvent at a controlled pH and the

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resulting organic phase is recycled to a suitable point in the styrene purification system.

The recycled solution is said to be a more effective inhibitor than the dinitrophenol originally employed.

U.S. Patent Number 3,966,711 teaches that 2,2,7,7-tetraalkyl- and 2,7-dispiroalkylene-5-oxo-1,4-diazacycloheptanes substituted in the 4-position by monoor tetravalent radicals are powerful light-stabilizers for organic polymers. They are said to possess higher compatibility than their 4-unsubstituted homologues, from which they can be synthesized by reactions known for N-alkylation. Preferred substituents in the 4-position are alkyl, alkylene, alkenyl, aralkyl, and esteralkyl groups. The 1-nitroxyls derived from the imidazolidines by oxidation with hydrogen peroxide or percarboxylic acids are also said to be good light stabilizers.

U.S. Patent Number 4,033,829 discloses the inhibition of styrene against polymerization during the distillation thereof by incorporating therein, in an amount sufficient to inhibit polymerization thereof, the dinitrophenol solution recovered from styrene still residues or tars resulting from the distillation of styrene in the presence of dinitrophenol. The recovered dinitrophenol solution is said to be a more effective polymerization inhibitor than dinitrophenol per se.

U.S. Patent Numbers 4,252,615 and 4,469,558 disclose a process for the distillation of readily polymerizable vinyl aromatic compounds and a polymerization inhibitor therefor. The process comprises subjecting a vinyl aromatic compound to elevated temperatures in a distillation system in the presence of a polymerization inhibitor comprising 2,6-dinitro-p-cresol. Also disclosed is a distillation method and apparatus for use with this inhibitor.

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U.S. Patent Number 4,665,185 discloses a process for the efficient preparation of nitroxyls of sterically hindered amines by the oxidation of the amine using a hydroperoxide in the presence of a small amount of a metal ion catalyst, at moderate temperature for a short period of time, to give the nitroxyl in high yield and purity.

U.S. Patent Number 4,774,374 discloses a vinyl aromatic composition stabilized against polymerization comprising (a) a vinyl aromatic compound and (b) an effective amount of a stabilizer system in which the active ingredient consists essentially of an oxygenated species formed by the reaction of oxygen and an N-aryl-N'-alkyl-p-phenylenediamine.

U.S. Patent Number 5,254,760 teaches that the polymerization of a vinyl aromatic compound, such as styrene, is very effectively inhibited during distillation or purification by the presence of at least one stable nitroxyl compound together with at least one aromatic nitro compound.

U.S. Patent Numbers 5,545,782 and 5,545,786 disclose that nitroxyl inhibitors in combination with some oxygen reduce the premature polymerization of vinyl aromatic monomers during the manufacturing processes for such monomers. Even small quantities of air used in combination with the nitroxyl inhibitors are said to result in vastly prolonged inhibition times for the monomers.

European Patent Application 0 178 168 A2 discloses a method for inhibiting the polymerization of an α,β -ethylenically unsaturated monocarboxylic acid during its recovery by distillation by using a nitroxide free radical.

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European Patent Application 0 765 856 A1 discloses a stabilized acrylic acid composition in which the polymerization of the acrylic acid is inhibited during the distillation process for purifying or separating the acrylic acid as well as during transport and storage. The compositions comprise three components: (a) acrylic acid, (b) a stable nitroxyl radical, and (c) a dihetero-substituted benzene compound having at least one transferable hydrogen (e.g., a quinone derivative such as the monomethyl ether of hydroquinone (MEHQ)). During the distillation process, transport, and storage, components (b) and (c) are present in a polymerization-inhibiting amount. During the distillation process, oxygen (d) is preferably added with components (b) and (c). According to the specification, examples of suitable nitroxide free radical compounds include di-t-butylnitroxide; di-t-amylnitroxide; 2,2,6,6-tetramethylpiperidinyloxy; 4-hydroxy-2,2,6,6-tetramethyl-piperidinyloxy; 4-oxo-2,2,6,6tetramethyl-piperidinyloxy; 4-dimethylamino-2,2,6,6-tetramethyl-piperidinyloxy; 4amino-2,2,6,6-tetramethyl-piperidinyloxy; 4-ethanoyloxy-2,2,6,6-tetramethylpiperidinyloxy; 2,2,5,5-tetramethylpyrrolidinyloxy; 3-amino-2,2,5,5tetramethylpyrrolidinyloxy; 2,2,5,5-tetramethyl-1-oxa-3-azacyclopentyl-3-oxy; 2,2,5,5tetramethyl-1-oxa-3-pyrrolinyl-1-oxy-3-carboxylic acid; and 2,2,3,3,5,5,6,6octamethyl-1,4-diazacyclohexyl-1,4-dioxy.

WO 97/46504 concerns substance mixtures containing: (A) monomers containing vinyl groups; and (B) an active amount of a mixture which inhibits premature polymerization of the monomers containing vinyl groups during their purification or distillation and contains: (i) between 0.05 and 4.5 wt %, relative to the total mixture (B), of at least one N-oxyl compound of a secondary amine which has no

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hydrogen atom at the α-C atoms; and (ii) between 99.95 and 95.5 wt % relative to the total mixture (B), of at least one nitro compound. The publication also discloses a process for inhibiting the premature polymerization of monomers, and the use of mixture (B) for inhibiting the premature polymerization of monomers.

WO 98/14416 discloses that the polymerization of vinyl aromatic monomers such as styrene is inhibited by the addition of a composition of a stable hindered nitroxyl radical and an oxime compound.

WO 98/25872 concerns substance mixtures containing: (A) compounds containing vinyl groups; (B) an active amount of a mixture which inhibits premature polymerization of the compounds containing vinyl groups and contains: (i) at least one N-oxyl compound of a secondary amine which does not carry any hydrogen atoms on the α-carbon atoms; and (ii) at least one iron compound; (C) optionally nitro compounds; and (D) optionally co-stabilizers. The publication also discloses a process for inhibiting the premature polymerization of compounds (A) containing vinyl groups, and the use of (B) optionally mixed with nitro compounds (C) and/or co-stabilizers (D) for inhibiting the premature polymerization of radically polymerizable compounds and stabilizing organic materials against the harmful effect of radicals.

U.K. Patent Number 1,127,127 discloses that acrylic acid can be stabilized against polymerization by the addition thereto of a nitroxide having the essential skeletal structure:

wherein R_1 , R_2 , R_3 , and R_4 are alkyl groups and no hydrogen is bound to the remaining valencies on the carbon atoms bound to the nitrogen. The two remaining valencies that are not satisfied by R_1 to R_4 or nitrogen can also form part of a ring (e.g., 2,2,6,6 tetramethyl-4-hydroxy-piperidine-1-oxyl).

UK Patent Application GB 2 069 523 A discloses carrying out the distillation of readily polymerizable vinyl aromatic compounds at elevated temperature in the presence of a polymerization inhibitor comprising 2,6-dinitro-p-cresol.

CS-260755 B1 is directed to the preparation of 4-substituted-2,2,6,6-tetramethylpiperidine nitroxyls as olefin stabilizers.

SU-334845 A1 is directed to the inhibition of the radical polymerization of oligoester acrylates using iminoxyl radical inhibitors of a given formula.

FR 2,761,060 relates to the prevention of premature polymerization of styrene during its production by dehydrogenation of ethylbenzene by injecting into the process effluent a radical inhibitor based on an oxyl-tetramethylpiperidine derivative.

The foregoing are incorporated herein by reference in their entirety.

SUMMARY OF THE INVENTION

During the production and purification of unsaturated monomers, many manufacturers recycle certain process streams containing the polymerization inhibitor in order to improve the economics of the process by reusing the inhibitor left in the process stream. It is known in the industry that the recycling of streams utilizing nitroxyls as polymerization inhibitors in plants employing temperatures in excess of about 115° C causes loss of inhibitor efficiency, such that the tar recycle leads to a higher polymer content than would be expected or desirable.

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It has now been discovered that recycling streams utilizing nitroxyls as polymerization inhibitors can be done effectively in some plants without loss of inhibitor efficiency.

More particularly, the present invention is directed to an improvement in a process for the production and purification of unsaturated monomers employing nitroxyl-containing inhibitors wherein process streams containing the inhibitor are recycled, wherein the improvement comprises recycling said streams at a reboiler temperature no higher than about 110° C.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

As stated above, the present invention is directed to inhibiting systems in which at least one nitroxyl radical is used in the purification train to prevent polymer growth that occurs via a "living" polymerization mechanism and, more particularly, to effectively recycling the nitroxyl-containing process streams at reboiler temperatures below about 110° C without loss of inhibitor efficiency.

In one preferred aspect, the inhibitor is a stable hindered nitroxyl compound having the structural formula:

wherein

 R_1 and R_4 are independently selected from the group consisting of hydrogen, alkyl, and heteroatom-substituted alkyl;

R₂ and R₃ are independently selected from the group consisting of alkyl and

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heteroatom-substituted alkyl; and

 X_1 and X_2

- (1) are independently selected from the group consisting of halogen, cyano, amido, -S-C₆H₅, carbonyl, alkenyl, alkyl of 1 to 15 carbon atoms, COOR₇, -S-COR₇, and -OCOR₇, wherein R₇ is alkyl or aryl, or
- (2) taken together, form a ring structure with the nitrogen.

In a particularly preferred embodiment, the stable hindered nitroxyl compound has the structural formula:

wherein R_1 and R_4 are independently selected from the group consisting of hydrogen, alkyl, and heteroatom-substituted alkyl and R_2 and R_3 are independently selected from the group consisting of alkyl and heteroatom-substituted alkyl, and the

portion represents the atoms necessary to form a five-, six-, or seven-membered heterocyclic ring.

Accordingly, one of the several classes of cyclic nitroxides that can be employed in the practice of the present invention can be represented by the following structural formula:

$$\begin{array}{c|c}
Z_1 \\
Z_2 \\
R_1 \\
R_2 \\
C^*
\end{array}$$

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wherein Z_1 , Z_2 , and Z_3 are independently selected from the group consisting of oxygen, sulfur, secondary amines, tertiary amines, phosphorus of various oxidation states, and substituted or unsubstituted carbon atoms, such as >CH $_2$, >CHCH $_3$, >C=O, >C(CH $_3$) $_2$, >CHBr, >CHCl, >CHI, >CHF, >CHOH, >CHON, >C(OH)CN, >CHCOOH, >CHCOOCH $_3$, >CHCOOC $_2$ H $_5$, >C(OH)COOC $_2$ H $_5$, >C(OH)COOC $_3$ H $_5$, >CENOH, >CENOH, >CENOH, >CENOH, >CECH-C $_6$ H $_5$, >CF $_2$, >CCI $_2$, >CBr $_2$, >CI $_3$, >CR $_5$ PR $_{13}$ R $_{14}$ R $_{15}$, and the like, where R $_5$ and R $_6$ are independently selected from the group consisting of hydrogen, alkyl, aryl, and acyl and R $_{13}$, R $_{14}$, and R $_{15}$ are independently selected from the group consisting of unshared electrons, alkyl, aryl, =O, OR $_{16}$, and NR $_{17}$ R $_{18}$, where R $_{16}$, R $_{17}$, and R $_{18}$ are independently selected from the group consisting of hydrogen, alkyl, and aryl. Where R $_5$ and/or R $_6$ are alkyl, it is preferred that they be a lower alkyl (i.e., one having one to five carbon atoms, e.g., methyl, ethyl, propyl, butyl, pentyl, and isomers thereof).

Where R₅ and/or R₆ are aryl, it is preferred that they be aryl of from 6 to 10 carbon atoms, e.g., phenyl or naphthyl, which, in addition, may be substituted with non-interfering substituents, e.g., lower alkyl groups, halogens, and the like.

Where R_5 and/or R_6 are acyl, it is preferred that they be acyl of the structure

where R_{19} is alkyl, aryl, OR_{20} , or $NR_{20}R_{21}$ and where R_{20} and R_{21} are alkyl, aryl, or

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where R_{22} is alkyl or aryl. Where R_{19} , R_{20} , R_{21} , or R_{22} are alkyl, they are preferably alkyl of from 1 to 15 carbon atoms, more preferably lower alkyl of from 1 to 5 carbon atoms, as described above. Where R_{19} , R_{20} , R_{21} , or R_{22} are aryl, they are preferably aryl of from 6 to 10 carbon atoms, as described above.

Another of the several classes of cyclic nitroxides that can be employed in the practice of the present invention can be represented by the following structural formula:

$$R_1$$
 R_2
 R_3
 R_3
 R_4

wherein Z_1 and Z_2 , which may be the same or different, are nitrogen or substituted or unsubstituted carbon atoms, such as =C(H)-, =C(CH₃)-, =C(COOH)-, =C(COOCH₃)-, =C(COOC₂H₅)-, =C(OH)-, =C(CN)-, =C(NR₅R₆)-, =C(CONR₅R₆)-, and the like, and where Z_3 , R_5 , and R_6 are as described above.

The cyclic nitroxides employed in the practice of the present invention can also be derived from five-membered rings. These compounds are of the structure:

$$R_1$$
 R_2
 R_3
 R_3

wherein Z₂ and Z₃, which may be the same or different, are sulfur, oxygen, secondary amines, tertiary amines, phosphorus of various oxidation states, or substituted or unsubstituted carbon atoms, such as, >CH₂, >CHCH₃, >C=O, >C(CH₃)₂, >CHBr, >CHCl, >CHI, >CHF, >CHOH, >CHCN, >C(OH)CN, >CHCOOH, >CHCOOCH₃, >CHCOOC₂H₅, >C(OH)COOC₂H₅, >C(OH)COOCH₃, >C(OH)CHOHC₂H₅, >CR₅OR₆, >CHNR₅R₆, >CCONR₅R₆, >C=NOH, >C=CH-C₆H₅, CF₂, CCl₂, CBr₂, Cl₂, >CR₅PR₁₃R₁₄R₁₅, and the like, wherein the several R groups are as described above.

The cyclic nitroxides employed in the practice of the present invention can also have the structure:

$$R_1$$
 R_2
 R_3
 R_3

wherein Z_4 and Z_5 , which can be the same or different, can be nitrogen or a substituted or unsubstituted carbon atom, such as =C(H)-, $=C(CH_3)$ -, =C(COOH)-, $=C(COOC_2H_5)$ -, =C(OH)-, =C(CN)-, $=C(NR_5R_6)$ -, $=C(CONR_5R_6)$ -, and the like, where R_5 and R_6 are as described above.

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Another class of cyclic nitroxides that can be employed in the practice of the present invention is of the structure:

$$R_2 \xrightarrow{R_1}^{N} \stackrel{0*}{\underset{Z_2 \cdot Z_3}{\bigvee}} R_3$$

wherein Z_2 and Z_3 , which may be the same or different, are sulfur, oxygen, secondary amines, tertiary amines, or substituted or unsubstituted carbon atoms, such as, >CH₂, >CHCH₃, >C=O, >C(CH₃)₂, >CHBr, >CHCl, >CHI, >CHF, >CHOH, >CHCN, >C(OH)CN, >CHCOOH, >CHCOOCH₃, >CHCOOC₂H₅, >C(OH)COOC₂H₅, >C(OH)COOCH₃, >C(OH)CHOHC₂H₅, >CHNR₅R₆, >CCONR₅R₆, >CR₅OR₆, >C=NOH, >C=CH-C₆H₅, CF₂, CCl₂, CBr₂, CI₂, >CR₅PR₁₃R₁₄R₁₅, and the like, where the several R groups are as described above.

Further, two or more nitroxyl groups can be present in the same molecule, for example, by being linked through one or more of the Z-type moieties by a linking group E, as disclosed in U.S. Patent Number 5,254,760, which is incorporated herein by reference.

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As stated above, for all the nitroxyl structures above, R₁ and R₄ are independently selected from the group consisting of hydrogen, alkyl, and heteroatom-substituted alkyl and R₂ and R₃ are independently selected from the group consisting of alkyl and heteroatom-substituted alkyl. The alkyl (or heteroatom-substituted alkyl) groups R₁ through R₄ can be the same or different and preferably contain 1 to 15 carbon atoms, e.g., methyl, ethyl, propyl, butyl, pentyl, hexyl, heptyl, octyl, nonyl, decyl, undecyl, dodecyl, tridecyl, tetradecyl, pentadecyl, and the like, and isomers thereof, e.g., t-butyl, 2-ethylhexyl, and the like. It is more preferred that R₁ through R₄ be independently selected from lower alkyls (or heteroatom-substituted lower alkyls) of one to five carbon atoms (e.g., methyl, ethyl, propyl, butyl, pentyl, and isomers thereof). Where heteroatom substituents are present, they can, for example, include halogen, oxygen, sulfur, nitrogen, and the like. It is most preferred that all of R₁ through R₄ be methyl.

Examples of suitable nitroxide free radical compounds that can be used in the practice of the present invention, include, but are not limited to:

N,N-di-tert-butylnitroxide;

N,N-di-tert-amylnitroxide;

N-tert-butyl-2-methyl-1-phenyl-propylnitroxide;

N-tert-butyl-1-diethylphosphono-2,2-dimethylpropylnitroxide;

2,2,6,6-tetramethyl-piperidinyloxy;

4-amino-2,2,6,6-tetramethyl-piperidinyloxy;

4-hydroxy-2,2,6,6-tetramethyl-piperidinyloxy;

4-oxo-2,2,6,6-tetramethyl-piperidinyloxy;

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- 4-dimethylamino-2,2,6,6-tetramethyl-piperidinyloxy;
- 4-ethanoyloxy-2,2,6,6-tetramethyl-piperidinyloxy;
- 2,2,5,5-tetramethylpyrrolidinyloxy;
- 3-amino-2,2,5,5-tetramethylpyrrolidinyloxy;
- 5 2,2,4,4-tetramethyl-1-oxa-3-azacyclopentyl-3-oxy;
 - 2,2,4,4-tetramethyl-1-oxa-3-pyrrolinyl-1-oxy-3-carboxylic acid;
 - 2,2,3,3,5,5,6,6-octamethyl-1,4-diazacyclohexyl-1,4-dioxy;
 - 4-bromo-2,2,6,6-tetramethyl-piperidinyloxy;
 - 4-chloro-2,2,6,6-tetramethyl-piperidinyloxy;
 - 4-iodo-2,2,6,6-tetramethyl-piperidinyloxy;
 - 4-fluoro-2,2,6,6-tetramethyl-piperidinyloxy;
 - 4-cyano-2,2,6,6-tetramethyl-piperidinyloxy;
 - 4-carboxy-2,2,6,6-tetramethyl-piperidinyloxy;
 - 4-carbomethoxy-2,2,6,6-tetramethyl-piperidinyloxy;
- 4-carbethoxy-2,2,6,6-tetramethyl-piperidinyloxy;
 - 4-cyano-4-hydroxy-2,2,6,6-tetramethyl-piperidinyloxy;
 - 4-methyl-2,2,6,6-tetramethyl-piperidinyloxy;
 - 4-carbethoxy-4-hydroxy-2,2,6,6-tetramethyl-piperidinyloxy;
 - 4-hydroxy-4-(1-hydroxypropyl)-2,2,6,6-tetramethyl-piperidinyloxy;
 - 4-methyl-2,2,6,6-tetramethyl-1,2,5,6-tetrahydropyridine -1-oxyl;
 - 4-carboxy-2,2,6,6-tetramethyl-1,2,5,6-tetrahydropyridine -1-oxyl;
 - 4-carbomethoxy-2,2,6,6-tetramethyl-1,2,5,6-tetrahydropyridine -1-oxyl;
 - 4-carbethoxy-2,2,6,6-tetramethyl-1,2,5,6-tetrahydropyridine -1-oxyl;

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- 4-amino-2,2,6,6-tetramethyl-1,2,5,6-tetrahydropyridine -1-oxyl;
 4-amido-2,2,6,6-tetramethyl-1,2,5,6-tetrahydropyridine -1-oxyl;
- 3,4-diketo-2,2,5,5-tetramethylpyrrolidinyloxy;
- 3-keto-4-oximino-2,2,5,5-tetramethylpyrrolidinyloxy;
- 5 3-keto-4-benzylidine-2,2,5,5-tetramethylpyrrolidinyloxy;
 - 3-keto-4,4-dibromo-2,2,5,5-tetramethylpyrrolidinyloxy;
 - 2,2,3,3,5,5-hexamethylpyrrolidinyloxy;
 - 3-carboximido-2,2,5,5-tetramethylpyrrolidinyloxy;
 - 3-oximino-2,2,5,5-tetramethylpyrrolidinyloxy;
 - 3-hydroxy-2,2,5,5-tetramethylpyrrolidinyloxy;
 - 3-cyano-3-hydroxy-2,2,5,5-tetramethylpyrrolidinyloxy;
 - 3-carbomethoxy-3-hydroxy-2,2,5,5-tetramethylpyrrolidinyloxy;
 - 3-carbethoxy-3-hydroxy-2,2,5,5-tetramethylpyrrolidinyloxy;
 - 2,2,5,5-tetramethyl-3-carboxamido-2,5-dihydropyrrole-1-oxyl;
 - 2,2,5,5-tetramethyl-3-amino-2,5-dihydropyrrole-1-oxyl;
 - 2,2,5,5-tetramethyl-3-carbethoxy-2,5-dihydropyrrole-1-oxyl;
 - 2,2,5,5-tetramethyl-3-cyano-2,5-dihydropyrrole-1-oxyl;
 - bis(1-oxyl-2,2,6,6-tetramethylpiperidin-4-yl)succinate;
 - bis(1-oxyl-2,2,6,6-tetramethylpiperidin-4-yl)adipate;
- bis(1-oxyl-2,2,6,6-tetramethylpiperidin-4-yl)sebacate;
 - bis(1-oxyl-2,2,6,6-tetramethylpiperidin-4-yl)n-butylmalonate;
 - bis(1-oxyl-2,2,6,6-tetramethylpiperidin-4-yl)phthalate;
 - bis(1-oxyl-2,2,6,6-tetramethylpiperidin-4-yl)isophthalate,

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bis(1-oxyl-2,2,6,6-tetramethylpiperidin-4-yl)terephthalate;

bis(1-oxyl-2,2,6,6-tetramethylpiperidin-4-yl)hexahydroterephthalate;

N,N'-bis(1-oxyl-2,2,6,6-tetramethylpiperidin-4-yl)adipamide;

N-(1-oxyl-2,2,6,6-tetramethylpiperidin-4-yl)-caprolactam;

N-(1-oxyl-2,2,6,6-tetramethylpiperidin-4-yl)-dodecylsuccinimide;

2,4,6-tris-[N-butyl-N-(1-oxyl-2,2,6,6-tetramethylpiperidin-4-yl)]-s-triazine;

4,4'-ethylenebis(1-oxyl-2,2,6,6-tetramethylpiperazin-3-one); and the like.

As used herein, the abbreviation TEMPO stands for 2,2,6,6-tetramethyl-1-piperidinyloxy. Thus, 4-amino-TEMPO is 4-amino-2,2,6,6-tetramethyl-1-piperidinyloxy; 4-hydroxy-TEMPO is 4-hydroxy-2,2,6,6-tetramethyl-1-piperidinyloxy (also known in the art as HTEMPO); 4-oxo-TEMPO is 4-oxo-2,2,6,6-tetramethyl-1-piperidinyloxy; and so on.

It is preferred that at least one nitroxide free radical compound employed in the practice of the present invention be 4-amino-TEMPO, 4-oxo-TEMPO, 4-hydroxy-TEMPO, or TEMPO.

Blends of two or more of the foregoing, e.g., 4-amino-TEMPO and 4-oxo-TEMPO, can also be employed.

Such stable nitroxide free radical compounds can be prepared by known methods. (See, for example, U.S. Patent Numbers 3,163,677; 3,334,103; 3,372,182; 3,422,144; 3,494,930; 3,502,692; 3,873,564; 3,966,711; and 4,665,185; which are incorporated herein by reference.) They are suitable for use over a wide range of temperatures, but distillation temperatures employed with the ethylenically unsaturated monomers that are stabilized with the nitroxide free radical compounds typically range

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from about 60° C to about 180° C, preferably from about 70° C to about 165° C, and, more preferably, from about 80° C to about 150° C. Such distillations are generally performed at an absolute pressure in the range of about 10 to about 1,200 mm of Hg.

The ethylenically unsaturated monomer, the premature polymerization and polymer growth of which is an object of the present invention, can be any such monomer for which unintended polymerization and/or polymer growth during its manufacture, storage, and/or distribution is a problem. Among those monomers that will benefit from the practice of the present invention are: styrene, α-methylstyrene, styrene sulfonic acid, vinyltoluene, divinylbenzenes, polyvinylbenzenes, alkylated styrene, 2-vinylpyridine, acrylonitrile, methacrylonitrile, methyl acrylate, ethyl acrylate, methyl methacrylate, ethyl methacrylate, acrylic acid, methacrylic acid, butadiene, chloroprene, isoprene, and the like

The ethylenically unsaturated monomers will not necessarily be stabilized indefinitely by the presence of the inhibitor(s), especially when the monomers are heated as in distillation, but they can be considered to be stabilized as long as A) there is a measurable increase in the time for which they can be heated before the onset of polymerization and/or polymer growth in a static system, B) the amount of polymer made at a constant temperature remains constant over time in a dynamic system, C) the rate of polymer growth is significantly slower than when the growth inhibiting system is not present, and/or D) the rate of polymer growth is nearly additive in the multi-pass test.

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Those skilled in the art will understand that, if desired, free radical scavengers can also be included in the stabilization process. For example, air or O₂, as disclosed in U.S. Patent Numbers 5,545,782 and 5,545,786, can be added, as can the aromatic nitro compounds disclosed in U.S. Patent Number 5,254,760, the dihetero-substituted benzene compounds having at least one transferable hydrogen, e.g., a quinone derivative such as the mono-methyl-ether of hydroquinone disclosed in European Patent Application 0 765 856 A1, the iron compounds disclosed in WO 98/25872, and other inhibitors, e.g., phenolics and certain inorganic salts, well-known to those skilled in the art.

The polymerization inhibitor(s) can be introduced into the monomer to be protected by any conventional method. They can, for example, be added as a concentrated solution in suitable solvents just upstream from the point of desired application by any suitable means. In addition, individual inhibiting components can be injected separately into the distillation train along with the incoming feed and/or through separate and multiple entry points, provided there is an efficient distribution of the inhibiting composition. Since the inhibitors are gradually depleted during the distillation operation, it is generally advantageous to maintain the appropriate amount of them in the distillation apparatus by adding them during the course of the distillation process. Adding inhibitors can be done either on a generally continuous basis or intermittently, in order to maintain the inhibitor concentration above the minimum required level.

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The total inhibitor concentration should be from about 1 to about 2000 ppm versus the monomer being inhibited; preferably from about 5 to about 1000 ppm, depending on the conditions of use.

The advantages and the important features of the present invention will be more apparent from the following examples.

EXAMPLES

Procedure for Multi-Pass Reboiler Test

Preparation of First Pass Feed Solution:

Tert-butylcatechol (TBC) is removed from commercially available styrene by distillation under vacuum. Removal of TBC is verified by caustic titration. The desired amount of inhibitor(s) is added to this TBC-free styrene either directly or by first making a concentrated solution of the inhibitor in TBC-free styrene followed by further dilution with TBC-free styrene.

Procedure for Reboiler Test (a dynamic test):

A quantity of the Feed Solution containing inhibitor (blend) at the desired charge (stated as a wt/wt total inhibitor to styrene) is added to a round-bottom flask (the Pot) and heated to the desired temperature (usually 130° C) and brought to reflux by adjusting the pressure/vacuum. Once the Pot contents are at temperature, a continuous stream of fresh Feed Solution is begun at a rate that will add the volume of the initial Pot solution to the Pot over a period of time called the residence time (typically, one hour). At the same time that the fresh Feed Solution flow is begun, the Bottoms Stream flow is also begun. The Bottoms Stream is solution in the Pot that is removed at the same rate as the fresh Feed Solution is added. The equal flows of Feed and Bottoms Streams cause the quantity in

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the Pot to remain constant over the time of the experiment, while allowing continuous replenishment of inhibitor. This procedure simulates the way inhibitors are used in a distillation train of a plant producing vinyl monomers. The experiment continues with flow in and out of the Pot for a specified period of time. Typically, the First Pass runs for 10 hours, the Second Pass runs for 9 hours, the Third Pass runs for 8 hours, etc.

Samples are collected hourly from the Bottoms Stream. These samples are analyzed for polymer content via the methanol turbidity method. Polymer Content is reported as average weight percent polymer (Ave. Wt % Polymer), which is the average of the polymer contents for the hourly samples after four hours running. The amount of polymer in the samples is an indication of effectiveness of the inhibitor system being tested.

Preparation of Second and Third Pass Feed Solutions:

The Bottoms Stream from the previous Pass is collected. The amounts of inhibitor(s) in the First Pass Feed Solution and the Bottoms Stream from the First Pass are determined by appropriate analytical method(s), e.g., gas chromatography. An amount of inhibitor(s) is added to the collective Bottoms Stream from the First Pass to increase the level of inhibitor(s) in the Bottoms Stream to a level equal to that found in the First Pass Feed Solution. For each subsequent Pass, an appropriate amount of inhibitor(s) is added to the collective Bottoms Streams to bring inhibitor(s) level equal to that found in the First Pass Feed Solution.

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Evaluation of the Results

The difference in the amount of polymer made in one Pass versus subsequent

Passes is an indication of the ability of the inhibiting system to prevent or allow

polymer to grow. For example, an increase in the amount of polymer made going from

one Pass to the next which is roughly equivalent to the amount of polymer made

during the First Pass is an indication that the inhibiting system effectively prevents

polymer growth during recycle. Conversely, an increase in the amount of polymer

made going from one Pass to the next that is dramatically greater (about 10 times or

more) than the amount of polymer made during the First Pass is an indication that the

inhibiting system does not effectively prevent polymer growth during recycle.

Recycle of Nitroxyl-containing Streams at Low Temperatures

The Multi-Pass test described above simulates recycle of styrene purification tar streams in a plant. As stated above, many companies recycle certain process streams containing the polymerization inhibitor in order to improve their economics by reusing the inhibitor left in the process stream. It has previously been noted that the recycle of streams utilizing nitroxyls as polymerization inhibitors in higher temperature (above 115° C) plants causes loss of inhibitor efficiency, such that recycle leads to higher polymer content than expected or desired. It has now been found that the recycle of streams utilizing nitroxyls as polymerization inhibitors in lower temperature plants (below about 110° C) can be done effectively without loss of inhibitor efficiency, as indicated by comparison of test results versus DNBP or a PDA/DNBP blend, both of which have been shown to be effectively utilized with tar recycle in low temperature and high temperature plants. The test results in Table 1 below illustrate

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the invention.

Table 1				
Inhibitor System	Temp (°C)	Ave. Wt % Polymer		
		Pass 1	Pass 2	Pass 3
1500 ppm PDA/DNBP Blend	130	0.066	0.183	0.423
1500 ppm DNBP	130	0.109	0.266	0.430
300 ppm Nitroxyl	130	0.052	1.45	7.45
100 ppm Nitroxyl	116	0.00051	0.0408	2.36
25 ppm Nitroxyl	100	0.00019	0.00022	0.00043

PDA is N-(1,4-dimethylpentyl)-N'-phenyl-p-phenylenediamine.

DNBP is 2,4-dinitro-6-sec-butylphenol.

Nitroxyl is 4-oxo-TEMPO.

The first two Examples indicate results for systems known in industry to recycle with excellent inhibitor efficiency. The third example reflects conditions known in industry to lead to loss of nitroxyl inhibitor efficiency. It will be noted in the above Table that as the temperature is lowered, not only does nitroxyl inhibitor efficiency improve in recycle, but also the amount of nitroxyl utilized can be lowered providing an additional advantage owing to the relatively high cost of the nitroxyl compounds.

In view of the many changes and modifications that can be made without departing from principles underlying the invention, reference should be made to the appended claims for an understanding of the scope of the protection to be afforded the invention.